10/019,436

=> file caplus

FILE 'CAPLUS' ENTERED AT 13:22:06 ON 16 JUL 2002

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FILE COVERS 1907 - 16 Jul 2002 VOL 137 ISS 3 FILE LAST UPDATED: 15 Jul 2002 (20020715/ED)

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=> d que L1

l STR

Structure attributes must be viewed using STN Express query preparation.

L3 2 SEA FILE=REGISTRY SSS FUL L1

L4 1 SEA FILE=CAPLUS L3

=> d l4 ibib abs hitstr

L4 ANSWER 1 OF 1 CAPLUS COPYRIGHT 2002 ACS ACCESSION NUMBER: 2001:31494 CAPLUS

appliants

DOCUMENT NUMBER:

134:86172

TITLE:

Preparation and effect of quinolinecarboxylic acid

derivative or salts as antibacterial agents

INVENTOR(S):

Yazaki, Akira; Niino, Yoshiko; Kuramoto, Yasuhiro;

Hirao, Yuzo; Oshita, Yoshihiro; Hayashi, Norihiro;

Amano, Hirotaka

PATENT ASSIGNEE(S):

Wakunaga Pharmaceutical Co., Ltd., Japan

SOURCE:

PCT Int. Appl., 25 pp. CODEN: PIXXD2

DOCUMENT TYPE:

Patent

LANGUAGE:

Japanese

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

PATENT NO. KIND DATE APPLICATION NO. DATE	DATE			
WO 2001002390 A1 20010111 WO 2000-JP4096 20000622	20000622			
W: AE, AG, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, CA, CH, CN,	CR,			
CU, CZ, DE, DK, DM, DZ, EE, ES, FI, GB, GD, GE, GH, GM, HR,	HU,			
ID, IL, IN, IS, JP, KE, KG, KP, KR, KZ, LC, LK, LR, LS, LT,	LU,			
LV, MA, MD, MG, MK, MN, MW, MX, MZ, NO, NZ, PL, PT, RO, RU,	SD,			
SE, SG, SI, SK, SL, TJ, TM, TR, TT, TZ, UA, UG, US, UZ, VN,	-			
ZA, ZW, AM, AZ, BY, KG, KZ, MD, RU, TJ, TM	•			
RW: GH, GM, KE, LS, MW, MZ, SD, SL, SZ, TZ, UG, ZW, AT, BE, CH,	CY,			
DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE, BF,	-			
CF, CG, CI, CM, GA, GN, GW, ML, MR, NE, SN, TD, TG	•			
EP 1193266 A1 20020403 EP 2000-940804 20000622				
R: AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, MC,	PT.			
IE, SI, LT, LV, FI, RO	,			
BR 2000012192 A 20020618 BR 2000-12192 20000622				
NO 2001006378 A 20020228 NO 2001-6378 20011227				
PRIORITY APPLN. INFO.: JP 1999-187492 A 19990701				
WO 2000-JP4096 W 20000622				
GI				

$$F$$
 CO_2H
 H_2N
 F

AB The title compd. I and salts were prepd. The title compd. I was characterized by, when orally administered, showing a long half-life in blood while sustaining an extremely high antibacterial effect and a low toxicity, and having an extremely high bioavailability. Thus, title compd. I is widely usable as preventives and remedies for various infectious diseases in humans and animals.

I

IT 318269-49-7P 318269-50-0P

RL: BAC (Biological activity or effector, except adverse); BSU (Biological study, unclassified); SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (Uses)

(prepn. and effect of quinolinecarboxylic acid deriv. or salts as antibacterial agents)

RN 318269-49-7 CAPLUS

CN 3-Quinolinecarboxylic acid, 1-(6-amino-3,5-difluoro-2-pyridinyl)-8-bromo-7-[3-(ethylamino)-1-azetidinyl]-6-fluoro-1,4-dihydro-4-oxo-(9CI) (CA INDEX NAME)

RN 318269-50-0 CAPLUS

CN 3-Quinolinecarboxylic acid, 1-(6-amino-3,5-difluoro-2-pyridinyl)-8-bromo-7-[3-(ethylamino)-1-azetidinyl]-6-fluoro-1,4-dihydro-4-oxo-, (2Z)-2-butenedioate (1:1) (9CI) (CA INDEX NAME)

CM 1

CRN 318269-49-7 CMF C20 H17 Br F3 N5 O3

CM 2

CRN 110-16-7 CMF C4 H4 O4 CDES 2:Z

Double bond geometry as shown.

REFERENCE COUNT:

THERE ARE 13 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

=> file caplus FILE 'CAPLUS' ENTERED AT 13:24:20 ON 16 JUL 2002 USE IS SUBJECT TO THE TERMS OF YOUR STN CUSTOMER AGREEMENT. PLEASE SEE "HELP USAGETERMS" FOR DETAILS. COPYRIGHT (C) 2002 AMERICAN CHEMICAL SOCIETY (ACS)

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FILE COVERS 1907 - 16 Jul 2002 VOL 137 ISS 3 FILE LAST UPDATED: 15 Jul 2002 (20020715/ED)

This file contains CAS Registry Numbers for easy and accurate substance identification.

CAS roles have been modified effective December 16, 2001. Please check your SDI profiles to see if they need to be revised. For information on CAS roles, enter HELP ROLES at an arrow prompt or use the CAS Roles thesaurus (/RL field) in this file.

=> d que

L5

STR

Structure attributes must be viewed using STN Express query preparation.

L6 3 SEA FILE=REGISTRY SSS FUL L5

L7 2 SEA FILE=CAPLUS L6

=> d 17 1-2 ibib abs hitstr

L7 ANSWER 1 OF 2 CAPLUS COPYRIGHT 2002 ACS

ACCESSION NUMBER:

2001:31494 CAPLUS

DOCUMENT NUMBER:

134:86172

TITLE:

Preparation and effect of quinolinecarboxylic acid

derivative or salts as antibacterial agents

INVENTOR(S):

Yazaki, Akira; Niino, Yoshiko; Kuramoto, Yasuhiro;

Hirao, Yuzo; Oshita, Yoshihiro; Hayashi, Norihiro;

Amano, Hirotaka

PATENT ASSIGNEE(S):

Wakunaga Pharmaceutical Co., Ltd., Japan

SOURCE: PCT Int. Appl., 25 pp.

CODEN: PIXXD2

DOCUMENT TYPE:

Patent

LANGUAGE:

Japanese

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PA	rent :	NO.		KI	ND	DATE			Α	PPLI	CATI	ON NO	ο.	DATE			
WO	0 2001002390			A1 20010111				WO 2000-JP4096 20000622									
	w:	ΑE,	AG,	AL,	AM,	AT,	AU,	ΑZ,	BA,	BB,	BG,	BR,	BY,	CA,	CH,	CN,	CR,
		CU,	CZ,	DE,	DK,	DM,	DZ,	EE,	ES,	FI,	GB,	GD,	GE,	GH,	GM,	HR,	HU,
		ID,	IL,	IN,	IS,	JP,	ΚE,	KG,	KP,	KR,	KZ,	LC,	LK,	LR,	LS,	LT,	LU,
		LV,	MA,	MD,	MG,	MK,	MN,	MW,	MX,	MZ,	NO,	NZ,	PL,	PT,	RO,	RU,	SD,
		SE,	SG,	SI,	SK,	SL,	ТJ,	TM,	TR,	TT,	TZ,	UA,	ŬĠ,	US,	UZ,	VN,	YU,
		ZA,	ZW,	AM,	ΑZ,	BY,	KG,	ΚZ,	MD,	RU,	ТJ,	TM					
	RW:	GH,	GM,	ΚE,	LS,	MW,	MZ,	SD,	SL,	SZ,	TZ,	UG,	ZW,	AT,	BE,	CH,	CY,
		DE,	DK,	ES,	FI,	FR,	GB,	GR,	IE,	IT,	LU,	MC,	NL,	PT,	SE,	BF,	ВJ,
			CG,			GΑ,											
ΕP	EP 1193266		A1 20020403				EP 2000-940804 2000062										
	R:	ΑT,	BE,	CH,	DE,	DK,	ES,	FR,	GB,	GR,	IT,	LI,	LU,	NL,	SE,	MC,	PT,
		ΙE,	SI,	LT,	LV,	FI,	RO										

20000622 BR 2000012192 Α 20020618 BR 2000-12192 20011227 20020228 NO 2001-6378 NO 2001006378 Α JP 1999-187492 19990701 PRIORITY APPLN. INFO .: Α WO 2000-JP4096 W 20000622

GΙ

AB The title compd. I and salts were prepd. The title compd. I was characterized by, when orally administered, showing a long half-life in blood while sustaining an extremely high antibacterial effect and a low toxicity, and having an extremely high bioavailability. Thus, title compd. I is widely usable as preventives and remedies for various infectious diseases in humans and animals.

Ι

IT 318269-49-7P 318269-50-0P

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(prepn. and effect of quinolinecarboxylic acid deriv. or salts as antibacterial agents)

RN 318269-49-7 CAPLUS

CN 3-Quinolinecarboxylic acid, 1-(6-amino-3,5-difluoro-2-pyridinyl)-8-bromo-7[3-(ethylamino)-1-azetidinyl]-6-fluoro-1,4-dihydro-4-oxo- (9CI) (CA INDEX NAME)

RN 318269-50-0 CAPLUS

CN 3-Quinolinecarboxylic acid, 1-(6-amino-3,5-difluoro-2-pyridinyl)-8-bromo-7-

[3-(ethylamino)-1-azetidinyl]-6-fluoro-1,4-dihydro-4-oxo-, (2Z)-2-butenedioate (1:1) (9CI) (CA INDEX NAME)

CM 1

CRN 318269-49-7

CMF C20 H17 Br F3 N5 O3

CM 2

CRN 110-16-7 CMF C4 H4 O4

CDES 2:Z

Double bond geometry as shown.

REFERENCE COUNT:

THERE ARE 13 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L7 ANSWER 2 OF 2 CAPLUS COPYRIGHT 2002 ACS

ACCESSION NUMBER:

1997:332364 CAPLUS

DOCUMENT NUMBER:

126:305587

TITLE:

Properation of novel pyridonecarboxylic acid

derivatives as antibacterial agents

INVENTOR(S):

Yazaki, Akira; Niino, Yoshiko; Ohshita, Yoshihiro;

Hirao, Yuzo; Amano, Hirotaka; Hayashi, Norihiro;

Kuramoto, Yasuhiro

PATENT ASSIGNEE(S):

Wakunaga Pharmaceutical Co., Ltd., Japan; Yazaki, Akira; Niino, Yoshiko; Ohshita, Yoshihiro; Hirao, Yuzo; Amano, Hirotaka; Hayashi, Norihiro; Kuramoto,

Yasuhiro

SOURCE:

PCT Int. Appl., 120 pp.

CODEN: PIXXD2

DOCUMENT TYPE:

Patent

LANGUAGE:

Japanese

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

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W: AU, BR, CA, CN, HU, JP, KR, MX, RU, US
             RW: AT, BE, CH, DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE
       CA 2232728 AA 19970327 CA 1996-2232728 19960920
AU 9670016 A1 19970409 AU 1996-70016 19960920
AU 707565 B2 19990715
CN 1201459 A 19981209 CN 1996-198104 19960920
EP 911327 A1 19990428 EP 1996-931264 19960920
       EP 911327 AI 199017 BI 20011205 DK ES.
             R: AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, MC, PT,
                   IE, FI
                                       19990727
       BR 9610485
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                                                              BR 1996-10485
                                                                                         19960920
                                       19991027
       EP 952151
                                A2
                                                             EP 1999-111114 19960920
                        A3 20010801
       EP 952151
            R: AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, MC, PT,
                   IE, FI
       JP 11322715
                                A2
                                       19991124
                                                              JP 1999-107671
                                                                                         19960920
                                                             EP 1999-124494 19960920
       EP 992501 A2 20000412
EP 992501 A3 20010905
                                         20000412
             R: AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, MC, PT,
   TE, FI

JP 3103380
B2 20001030
JP 1997-512602 19960920
RU 2167873
C2 20010527
RU 1998-107259 19960920
RU 2171252
C2 20010727
RU 1999-117921 19960920
AT 210129
E 20011215
AT 1996-931264 19960920

US 5998436
A 19991207
US 1998-43472 19980320
AU 9931227
Al 19990812
AU 721805
B2 20000713

US 6133284
A 20001017
US 1999-329246 19990610
US 6156903
AU 727457
B2 20001214
AU 79950144
AU 9950144
AI 20000330
JP 2000136191
A2 20000516
JP 3187795
B2 20010711
CN 1258672
A 20000705
CN 1999-123310 19991022
                   IE, FI
                               A 20000705
A 20000705
                                                           CN 1999-123310 19991022
CN 1999-123311 19991022
       CN 1258672
       CN 1258674
                                                           JP 1995-269280 A 19950922
JP 1996-178462 A 19960619
PRIORITY APPLN. INFO.:
                                                           AU 1996-70016 A3 19960920
EP 1996-931264 A3 19960920
                                                           JP 1997-512602 A3 19960920
                                                           WO 1996-JP2710 W 19960920
OTHER SOURCE(S): MARPAT 126:305587
GΙ
```

$$R^{5}$$
 R^{6}
 $Co_{2}R^{1}$
 N
 X
 Z
 R^{2}
 R^{3}

Ι

AB The title compds. (I; R1 = H, carboxyl protecting group; R2 = OH, lower alkoxy, optionally substituted amino; R3, R4 = H, halo; R5 = optionally halogenated or substituted satd. cyclic amino; R6 = H, halo, NO2, optionally protected amino; X, Y, and Z = N, :CH, :CR7; R7 = lower alkyl, halo, cyano, provided that at least one of X, Y, and Z = N; W = N, :CR8; R8 = H, halo, lower alkyl) are prepd. I are useful as antibacterial, antiviral, and anti-HIV agents. Thus, quinoline deriv. (II; X = F) was reacted with 3-aminoazetidine.2HCl in the presence of N-methylpyrrolidine to give the title compd. II (X = 3-aminoazetidin-1-yl), which showed MIC of < 0.013, 0.025, and 0.05 .mu.g/mL antibacterial effect for S. aureus, S. epidermidis, and P. aeruginosa resp.

IT 189280-18-0P

RL: BAC (Biological activity or effector, except adverse); BSU (Biological study, unclassified); SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (Uses)

(properation of novel pyridonecarboxylic acid derivs. as antibacterial agents)

RN 189280-18-0 CAPLUS

CN 3-Quinolinecarboxylic acid, 1-(6-amino-3,5-difluoro-2-pyridinyl)-8-chloro-7-[3-(ethylamino)-1-azetidinyl]-6-fluoro-1,4-dihydro-4-oxo- (9CI) (CA INDEX NAME)

=> file uspatall
FILE 'USPATFULL' ENTERED AT 13:24:59 ON 16 JUL 2002
CA INDEXING COPYRIGHT (C) 2002 AMERICAN CHEMICAL SOCIETY (ACS)

FILE 'USPAT2' ENTERED AT 13:24:59 ON 16 JUL 2002 CA INDEXING COPYRIGHT (C) 2002 AMERICAN CHEMICAL SOCIETY (ACS)

=> s 16

L8 3 L6

=> d 18 1-3 ibib abs hitstr

ANSWER 1 OF 3 USPATFULL

ACCESSION NUMBER: 2000:164659 USPATFULL

TITLE: Pyridonecarboxylic acid derivatives or their salts, and

antibacterial agents containing the same as their

effective components

INVENTOR(S): Yazaki, Akira, Takata-gun, Japan

Niino, Yoshiko, Takata-gun, Japan Ohshita, Yoshihiro, Takata-gun, Japan

Hirao, Yuzo, Takata-gun, Japan Amano, Hirotaka, Takata-gun, Japan Hayashi, Norihiro, Takata-gun, Japan Kuramoto, Yasuhiro, Takata-gun, Japan

PATENT ASSIGNEE(S): Wakunaga Pharmaceutical Co., Ltd., Osaka, Japan

(non-U.S. corporation)

NUMBER KIND DATE ______

PATENT INFORMATION: US 6156903 20001205 APPLICATION INFO.: US 1999-329336 19990610 (9)

RELATED APPLN. INFO.: Division of Ser. No. US 43472

NUMBER DATE ______

JP 1995-269280 19950922 PRIORITY INFORMATION:

JP 1996-178462 19960619

DOCUMENT TYPE: Utility · FILE SEGMENT:

Granted Seaman, D. Margaret PRIMARY EXAMINER:

NUMBER OF CLAIMS: EXEMPLARY CLAIM: LINE COUNT: 3385

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

A pyridonecarboxylic acid derivative represented by the following general formula (1): ##STR1## [wherein R.sup.1 represents hydrogen atom or a carboxyl protective group; R.sup.2 represents hydroxyl group, a lower alkoxy group, or a substituted or unsubstituted amino group; R.sup.3 represents hydrogen atom or a halogen atom; R.sup.4 represents hydrogen atom or a halogen atom; R.sup.5 represents a halogen atom or an optionally substituted saturated cyclic amino group; R.sup.6 represents hydrogen atom, a halogen atom, nitro group, or an optionally protected amino group; X, Y and Z may be the same or different and respectively represent nitrogen atom, --CH.dbd. or --CR.sup.7 .dbd. (wherein R.sup.7 represents a lower alkyl group, a halogen atom, or cyano group) (with the proviso that at least one of X, Y and Z represent the nitrogen atom), and W represents nitrogen atom or -- CR. sup.8 .dbd. (wherein R.sup.8 represents hydrogen atom, a halogen atom, or a lower alkyl group)] or its salt, as well as an antibacterial agent containing such compound are provided.

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

IT 189280-18-0P

(properation of novel pyridonecarboxylic acid derivs. as antibacterial agents)

189280-18-0 USPATFULL RN

3-Quinolinecarboxylic acid, 1-(6-amino-3,5-difluoro-2-pyridinyl)-8-chloro-CN 7-[3-(ethylamino)-1-azetidinyl]-6-fluoro-1,4-dihydro-4-oxo- (9CI) INDEX NAME)

ANSWER 2 OF 3 USPATFULL

ACCESSION NUMBER:

2000:138370 USPATFULL

TITLE:

Pyridonecarboxylic acid derivatives or their salts, and

antibacterial agents containing the same as their

effective components

INVENTOR(S):

Yazaki, Akira, Hiroshima-ken, Japan Niino, Yoshiko, Hiroshima-ken, Japan Ohshita, Yoshihiro, Hiroshima-ken, Japan

Hirao, Yuzo, Hiroshima-ken, Japan Amano, Hirotaka, Hiroshima-ken, Japan Hayashi, Norihiro, Hiroshima-ken, Japan Kuramoto, Yasuhiro, Hiroshima-ken, Japan

PATENT ASSIGNEE(S):

Wakunaga Pharmaceutical Co., Ltd., Osaka, Japan

(non-U.S. corporation)

NUMBER KIND DATE

PATENT INFORMATION: APPLICATION INFO.:

US 6133284 20001017 US 1999-329246 19990610 (9)

RELATED APPLN. INFO.: Division of Ser. No. US 43472

> NUMBER DATE -----

PRIORITY INFORMATION:

JP 1995-269280 19950922 JP 1996-178462

DOCUMENT TYPE:

Utility Granted

FILE SEGMENT:

PRIMARY EXAMINER:

Seaman, D. Margaret

LEGAL REPRESENTATIVE:

Birch, Stewart, Kolasch & Birch, LLP

NUMBER OF CLAIMS: EXEMPLARY CLAIM: 1 LINE COUNT: 3267

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

AΒ A pyridonecarboxylic acid derivative represented by the following general formula (1): ##STR1## [wherein R.sup.1 represents hydrogen atom or a carboxyl protective group; R.sup.2 represents hydroxyl group, a lower alkoxy group, or a substituted or unsubstituted amino group; R.sup.3 represents hydrogen atom or a halogen atom; R.sup.4 represents hydrogen atom or a halogen atom; R.sup.5 represents a halogen atom or an optionally substituted saturated cyclic amino group; R.sup.6 represents hydrogen atom, a halogen atom, nitro group, or an optionally protected amino group; X, Y and Z may be the same or different and respectively represent nitrogen atom, --CH.dbd. or --CR.sup.7 .dbd. (wherein R.sup.7 represents a lower alkyl group, a halogen atom, or cyano group) (with the proviso that at least one of X, Y and Z represent the nitrogen atom), and W represents nitrogen atom or --CR.sup.8 .dbd. (wherein R.sup.8 represents hydrogen atom, a halogen atom, or a lower alkyl group)]

or its salt, as well as an antibacterial agent containing such compound are provided.

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

IT 189280-18-0P

(properation of novel pyridonecarboxylic acid derivs. as antibacterial agents)

RN 189280-18-0 USPATFULL

CN 3-Quinolinecarboxylic acid, 1-(6-amino-3,5-difluoro-2-pyridinyl)-8-chloro-7-[3-(ethylamino)-1-azetidinyl]-6-fluoro-1,4-dihydro-4-oxo-(9CI) (CA INDEX NAME)

L8 ANSWER 3 OF 3 USPATFULL

ACCESSION NUMBER:

1999:160052 USPATFULL

TITLE:

Pyridonecarboxylic acid derivatives or their salts and antibacterial agent comprising the same as the active

ingredient

INVENTOR(S):

Yazaki, Akira, Hiroshima-ken, Japan Niino, Yoshiko, Hiroshima-ken, Japan Ohshita, Yoshihiro, Hiroshima-ken, Japan Hirao, Yuzo, Hiroshima-ken, Japan

Hirao, Yuzo, Hiroshima-ken, Japan Amano, Hirotaka, Hiroshima-ken, Japan Hayashi, Norihiro, Hiroshima-ken, Japan Kuramoto, Yasuhiro, Hiroshima-ken, Japan

PATENT ASSIGNEE(S):

Wakunaga Pharmaceuticals Co., Ltd., Osaka, Japan

(non-U.S. corporation)

	NUMBER	KIND DATE	
PATENT INFORMATION:	US 5998436 WO 9711068	19991207 19970327	
APPLICATION INFO.:	US 1998-43472 WO 1996-JP2710	19980320 19960920 19980320	(9) PCT 371 date

19980320 PCT 102(e) date

NUMBER DATE
-----PRIORITY INFORMATION: JP 1995-269280 19950922
JP 1996-178462 19960619

DOCUMENT TYPE: Utility FILE SEGMENT: Granted

PRIMARY EXAMINER: Mach, D. Margaret

LEGAL REPRESENTATIVE: Birch, Stewart, Kolasch & Birch, LLP

NUMBER OF CLAIMS: 11
EXEMPLARY CLAIM: 1
LINE COUNT: 3409

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

AΒ A pyridonecarboxylic acid derivative represented by the following general formula (1): ##STR1## [wherein R.sup.1 represents hydrogen atom or a carboxyl protective group; R.sup.2 represents hydroxyl group, a lower alkoxy group, or a substituted or unsubstituted amino group; R.sup.3 represents hydrogen atom or a halogen atom; R.sup.4 represents hydrogen atom or a halogen atom; R.sup.5 represents a halogen atom or an optionally substituted saturated cyclic amino group; R.sup.6 represents hydrogen atom, a halogen atom, nitro group, or an optionally protected amino group; X, Y and Z may be the same or different and respectively represent nitrogen atom, --CH.dbd. or --CR.sup.7 .dbd. (wherein R.sup.7 represents a lower alkyl group, a halogen atom, or cyano group) (with the proviso that at least one of X, Y and Z represent the nitrogen atom), and W represents nitrogen atom or -- CR. sup. 8 .dbd. (wherein R.sup.8 represents hydrogen atom, a halogen atom, or a lower alkyl group)] or its salt, as well as an antibacterial agent containing such compound are provided.

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